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## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:

Francis T. GILES et al.

Serial No.: 10/729,387

Group Art Unit: 1614

Filed: December 8, 2003

Examiner: Unassigned

For: PHARMACEUTICAL COMBINATIONS AND METHODS FOR THE TREATMENT OF

LEUKEMIA

## **INFORMATION DISCLOSURE STATEMENT UNDER 37 CFR §§ 1.56, 1.97 and 1.98**

Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

Sir:

This information disclosure statement is made in accordance with 37 C.F.R.  $\S\S$  1.56, 1.97 and 1.98 as follows:

## Timing and Fees

$\boxtimes$	Under 37 C.F.R. § 1.97(b), no fee or statement is required for filing this information disclosure statement is filed:						
		within three months of the filing date of a national application other than under § 1.53(d);					
			three months of the actual filing date of the national phase of a PCT ation; OR				
	$\boxtimes$	before the mailing of a first substantive office action (including after filing RCE).					
	Under 37 C.F.R. § 1.97(c), this information disclosure statement is filed after the perispecified in 37 C.F.R. § 1.97(b), but before the mailing date of:						
			a final rejection under 37 C.F.R. 1.113;				
			termination of prosecution, e.g. Ex Parte Quayle, M.P.E.P § 609(B)(2); OR				
			a notice of allowance under 37 C.F.R. § 1.311; and				

		is acco	mpanied by:
			the statement as specified in 37 C.F.R. § 1.97(e) set out below; OR
•			a check covering the fee of \$180.00 under 37 C.F.R. § 1.17(p).
			R. § 1.97(d), this information disclosure statement is filed after the mailing owing actions which have not been withdrawn:
			a final action under 37 C.F.R. § 1.113;
			termination of prosecution, e.g. Ex Parte Quayle, M.P.E.P $\S$ 609(B)(2); OR
			a notice of allowance under 37 C.F.R. § 1.311;
	AND i	s filed o	on or before payment of the issue fee; AND is accompanied by:
	·		the statement as specified in 37 C.F.R. § 1.97(e) as set forth below, and the fee of \$180.00 under 37 C.F.R. § 1.17(p).
Statem	ients Ui	nder 37	C.F.R. 1.97(e)
			Each item of information contained in this information disclosure statement was a first cited in a communication from a foreign patent office in a counterpart foreign application having a mailing date not more than three months prior to the filing date of this information disclosure statement; or
			No item of information contained in this information disclosure statement was cited in a communication from a foreign patent office in a counterpart foreign application, and to the knowledge of the undersigned attorney after making reasonable inquiry, no item of information contained in this information disclosure statement was known to any individual designated in 37 C.F.R. § 1.56(c) more than three months prior to the filing date of the information disclosure statement.
Cited ]	<u>Materia</u>	<u>ls</u>	
=0		ancest	of materials listed but not attached were cited in benefit (35 U.S.C. § 120) or application Serial No, on Form 892 by the Examiner and/or Form by the applicant; see 37 C.F.R. § 1.98(d).
		Copies 26, 20	s of materials listed were cited in an international search report dated <u>March</u> 04.

	$\boxtimes$	Copies of the materials listed are attached (except for the foregoing).
Non-E	English I	Language References
	$\boxtimes$	An English-language search report or equivalent paper from a foreign patent office is provided indicating the relevance of the cited reference(s).
		A foreign-language search report from a foreign patent office is provided, and pertinent parts are translated substantively below:
		<ul> <li>X = document of particular relevance when it is taken alone</li> <li>Y = document of particular relevance when it is combined with another such document</li> </ul>
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		<ul> <li>P = intercalated document</li> <li>T = document cited to understand the theory or principle underlying the invention</li> </ul>
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	A chec	ck for \$ covering the fee identified above is attached.
	Please	charge to Deposit Account No. 13-3402 \$ for the fee identified above.

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Respectfully submitted,

Brion P. Heaney, Reg. No. 32,542 Attorney/Agent for Applicants

MILLEN, WHITE, ZELANO & BRANIGAN, P.C.
Arlington Courthouse Plaza 1
2200 Clarendon Blvd. Suite 1400
Arlington, Virginia 22201
Telephone: (703) 243-6333
Facsimile: (703) 243-6410

Attorney Docket No. PHARMA-139

Date: June 21, 2004

BPH/rrt

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				Application Number	10/729,387			
INFO	RMATION	DIS	CLOSURE	Filing Date	December 3, 2003			
STATEMENT BY APPLICANT.				First Named Inventor	Francis T. GILES			
• . , .				Group Art Unit	1614			
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Sheet	1	of	2	Attorney Docket Number	PHARMA-139			

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## INFORMATION DISCLOSURE STATEMENT BY APPLICANT

Complete if Known 10/729,387 **Application Number** December 8, 2003 Filing Date Francis T. GILES First Named Inventor 1614 Group Art Unit **Examiner Name** Unassigned PHARMA-139 Attorney Docket Number

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OTHER PRIOR ART -- NON PATENT LITERATURE DOCUMENTS Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of Cite the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue Examiner No. number(s), publisher, city and/or country where published. Initials 1 Orsolic, Nada et al. "Troxatyl and STI571 Combination Therapy for Chronic Myeloid Leukemia: Preclinical In Vitro and In Vivo Evaluation", Blood, vol. 100, no. 11, (2002), Abstract No. 3107, XP0009027132 "FDA Approves Gleevec for Leukemia Treatment" FDA Consumer, US Dept. of Health, Educ. And Welfare, Public Health Services, US, vol. 4, July 2001, pg. 6, XP001145627 Giles F. et al. "Phase II Study of Troxatyl in Patients with Chronic Myeloid Leukemia in Blastic Phase (CML-BP)" Blood, W.B. Saunders Company, Orlando, FL, US, vol. 98, no. 11, part 2, (2001), pg. 258B, XP009007253

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